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- Method for detecting metals.
- Metals such as Ca²⁺, Sr²⁺, Sm³⁺, Cd²⁺ and lanthanoids such as La³⁺, Tb³⁺ and Yb³⁺ are detected by measuring the luminescence produced in contact with reproduced aequorin obtained by adding coelenterazine or its analogues to an enzyme of aequorin (apoaequorin).

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METHOD FOR DETECTING METALS

This invention relates to a method for detecting metals and especially to such a method in which a photoprotein is used, namely an enzyme of aequorin (apoaequorin), together with a substrate, as an emitter, namely coelenterazine or its analogues.

Aequorin is a photoprotein obtainable from luminescent jellyfish which grow in the northern part of the west coast of America. When one molecule of aequorin is specifically bound to two or three molecules of Ca²⁺, it oxidises the coelenterazine present, which acts as an emitter to produce luminescence. Thus, natural aequorin exists in the package state containing the enzyme

15 (apoaequorin), the substrate (coelenterazine) and molecular

oxygen (O₂). When Ca²⁺ is added to natural aequorin, its reaction with the Ca²⁺-binding sites present causes the coelenterazine to be oxidised, so that oxycoelenterazine, CO₂ and hv (emission) are produced. This light emission can be detected by a photomultiplier and its sensitivity is so high that trace concentrations of Ca²⁺ of about 10⁻⁷M are measurable.

The habitat of the jellyfish which produce natural aequorin is limited to the aforesaid coastal region and the season in which they are produced is also

limited. The yield of natural aequorin is only about 10 mg per 50,000 jellyfish. It is thus very important to obtain aequorin, but a certain supply cannot be assured.

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The inventor of the present invention has cloned the cDNA for aequorin obtained from these jellyfish by a recombinant DNA method. This cDNA clone was named plasmid pAQ440 (Japanese Patent Application 59-176125). Then, a plasmid containing a promoter in which the pAQ440 gene was inserted was transformed in a bacterium and it was found that the natural and fused type of aequorin (apoaequorin) could be efficiently expressed in <u>E.coli</u> (Japanese Patent Application 60-280259). This method provided a stable supply of aequorin.

The present invention further improves the method for detecting ${\rm Ca}^{2+}$ concentrations using natural aequorin and also provides a method for detecting metals, such as ${\rm Ca}^{2+}$ and others, using apoaequorin biosynthesised in E.coli.

According to one aspect of the present invention, a method for detecting metals is provided, wherein an enzyne of aequorin (apoaequorin) produced by biosynthesis in E.coli is used and wherein a substrate employed is coelenterazine or its analogues.

According to another aspect of this invention, a recombinant DNA molecule is characterised by comprising biologically pure aequorin synthesised by reacting coelenterazine or its analogues with apoaequorin.

Preferably, in carrying out the method of the invention, the aequorin enzyme (apoaequorin) is produced using the aequorin cDNA obtained by the biosynthesis method disclosed in the above-mentioned patent application and then, by adding coelenterazine as a substrate to the enzyme, aequorin having the same properties as natural aequorin is obtained. The luminescence shown by various metals when reacted with this synthesised aequorin is

measured, in order to detect and/or measure the amount of such metals. As a result, the presence of Ca²⁺, Sr²⁺ and Sm³⁺ can be observed. For Cd²⁺, its presence can be confirmed from the inhibitory rate of luminescence of Ca²⁺. Lanthanoids such as La³⁺, Tb³⁺ and Yb³⁺ also can be detected.

The following description serves to illustrate the invention more specifically.

(1) A method for producing aequorin (apoaequorin) in E.coli.

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(1) Insertion of aequorin genes into an expression vector having a promoter.

As the expression vector into which the aequorin gene is inserted, the vector in which the promoter is cloned is used. As the expression vector, pUC9 can be cited as an example. As the promoter, <u>lac</u>, <u>tac</u> or <u>trp</u> derived from <u>E.coli</u> or PL of the λ phage can be used, by way of example.

Firstly, pDR540 having the promoter is digested

20 by means of the restriction enzyme BamHI-HindIII. The
resulting fragment containing the promoter is then separated
and extracted by an electrophoresis method, e.g. in 8%
acrylamide gel. By using the promoter thus extracted, the
expression vector (piC9) can be obtained by cloning at the

25 BamHI-HindIII site of the vector.

Then, after digesting this expression vector with $\underline{\text{HindIII}}$, its terminals are repaired by $\underline{\text{E.coli}}$ DNA polymerase (Klenow fragment) in the presence of dATP, dGTP, dCTP and dTTP, both the terminals are ligated with \mathbf{T}_4 ligase and an expression vector, piC10, is thus obtained.

In preparing to bind the promoter in the piC10 with the aequorin cDNA gene, a synthetic nucleotide as a linker is cloned to piC10. Thus, the synthetic nucleotide linkers AR(5'GATCGATGGTCA-3') and AQ(5'AGCTTGACCATC-3') are first prepared, by a known synthetic method, and then annealed, after which phosphorylation of their 5'-terminals

is carried out using T₄ nucleotide kinase in the presence of ATP. The nucleotides thus phosphorylated are subsequently cloned, so as to construct plasmids in the form of repetition units of the linker at the BamHI-EcoRI site of the piC10, which has been constructed as mentioned above, and a recombinant DNA piC11 is so obtained.

Next, a <u>HindIII-Eco</u>RI fragment is separated from aequorin cDNA clone pAQ440 (see Japanese Patent Application 59-176125), this fragment is then inserted into the

- HindIII-EcoRI site of the above piC11, and the desired piQ5, namely bacteria containing a promoter in which the apoaequorin gene has been transformed, is obtained. This piQ5 has the ability to produce aequorin proteins of natural type.
- (2) Insertion of aequorin genes into an expression vector (pUC9, 9-1, 9-2) having a lac promoter in order to produce aequorin proteins of a fused type.

In a similar manner to that described in (1) above, PstI-EcoRI and HindIII-EcoRI fragments are separated from the cDNA clone pAQ440 and purified. Further, expression vectors pUC9, 9-1, 9-2 having a Lac promoter prepared by a similar method to that described in (1) above are obtained. Each of these fragments is cloned at the restriction enzyme site of each of these vectors to

- produce piQ9PE, 9-1PE, 9-2PE, 9-HE and 9-2HE. All of these piQ plasmids are under the control of the <u>lac</u> promoter and thus comprise aequorin proteins of the fused type having 8-amino acid residues at an N-terminus.
- (3) Production of proteins having aequorin activity using 30 <u>E.coli</u>.

Each plasmid cloned to the expression vectors obtained by the methods of (1) and (2) is transformed in a strain, such as $\underline{E.coli}$ (D 1210), and aequorin proteins are produced. In this method, for example, $\underline{E.coli}$ strains

35 containing plasmids are added to a certain volume of LB

broth containing a given concentration of ampicillin and the <u>E.coli</u> strains are cultured. Subsequently, an expression-inducible reagent is added to the culture medium and incubation is continued. The resulting culture medium is separated by centrifugation and the resulting cells are collected and washed with water.

(2) A method for synthesising coelenterazine and its analogues.

In this method, coelenterazine obtained by a known method can be used.

The coelenterazine used here has the following general formula:

$$\begin{array}{c|c}
0 & R_1 \\
 & N & N \\
R_3 & H
\end{array}$$

Coelenterazine of the natural type is represented by the formula wherein R₁ is $p-CH_2C_6H_4OH$, R₂ is $-C_6H_5$ and R₃ is $p-C_6H_4OH$.

Coelenterazine compounds represented by the formula (I) can be obtained by a known method, for example, the method disclosed by Inoue et al. (Published by Japan Chemical Society, Chemistry Letters, 141-144, 1975).

Further, as analogues of coelenterazine, analogue 1, represented by the formula (I) wherein R_1 is $-CH_2C_6H_5$, R_2 is $-C_6H_5$ and R_3 is $p-C_6H_4OH$, or analogue 2, represented by the formula (I) wherein R_1 is $-CH_3$, R_2 is $-C_6H_5$ and R_3 is $p-C_6H_4OH$, can be used. These analogues can be obtained by a known method, for example, the method disclosed by Halt et al. (Biochemistry, 18, (11) 2204-2210, 1979).

Metals are detected by the following method, for

35 example.

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The enzyme solution obtained by extraction from

E.coli using the above process (3) is used. After dissolving the enzyme solution in Tris-HCl buffer solution containing EDTA, coelenterazine as a substrate and 2
mercaptoethanol are added to the solution and it is then allowed to stand on ice, while aequorin is reproduced. A sample to be analysed is poured into the solution containing aequorin thus reproduced, and the resulting mixed solution is transferred to the reaction cell of a spectrophotometer. A sample to be measured is then injected into the cell and the quantity of luminescence produced is measured.

By using this invention, metals can be detected by means of aequorin produced in <u>E.coli</u> by the recombinant DNA method, using coelenterazine or its analogues as the substrate. Therefore, without using natural aequorin, which is difficult to obtain, metals can be detected by readily available synthetic aequorin produced in <u>E.coli</u>, and so can be easily, efficiently and economically detected.

The following non-limitative examples illustrate this invention more specifically.

EXAMPLE 1

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Production of aequorin (apoaequorin) from E.coli Construction of piC10:

The plasmid pDR540 having a tac promoter (produced by PL Pharmacia Co. Ltd.) was digested by the restriction enzyme BamHI-HindIII and then the resulting fragment containing the tac promoter of 92bp was separated and extracted by the electrophoresis method using acrylamide. Then, the BamHI-HindIII site of the resultant vector pUC9 was cloned and piC9 was constructed. Then, in order to remove the HindIII site of the expression vector piC9, firstly the HindIII was digested and then, after repairing the terminals of the vector by E.coli DNA polymerase (Klenow fragment) in the presence of dATP, dCTP and dTTP, both the terminals were ligated with

1 T₄ ligase and piC10 was thus constructed. EXAMPLE 2

Construction of piC11:

In order to bind the aequorin cDNA gene and the promoter, synthetic linker nucleotides of AR(5'GATCGATGGTCA-3!) and AQ(5'AGCTTGACCATC-3') were prepared. After these nucleotides were annealed, the 5' terminals were phosphorylated with T₄ nucleotide kinase in the presence of ATP. The nucleotides thus phosphorylated were subsequently cloned, so as to construct in the form of repetition units of the linker at the BamHI-EcoRI site of the piC10 (EXAMPLE 1), and piC11 was constructed.

EXAMPLE 3

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Construction of piQ5:

The <u>HindIII-Eco</u>RI fragment was separated and prepared from aequorin cDNA clone pAQ440, the fragment was inserted into the <u>HindIII-Eco</u>RI site of the above piC11, and piQ5 was thus obtained. This piQ5 can reproduce aequorin protein of the natural type.

20 EXAMPLE 4

Construction of piQ9PE, 9-1PE, 9-2PE, 9-HE and 9-2HE

After PstI-EcoRI and HindIII-EcoRI fragments

were separated and purified from cDNA clone pAQ440, these fragments were cloned at the restriction enzyme site of expression vectors pUC9, 9-1, 9-2, having a <u>lac</u> promoter, and piQ9PE, 9-1PE, 9-2PE, 9-HE and 9-2HE were constructed. All of these piQ fragments were under the control of the <u>lac</u> promoter and so comprised aequorin proteins of the fused type having 8-amino-acid residues at an N-terminus.

30 EXAMPLE 5

Production of proteins having aequorin activity using E.coli:

The above <u>E.coli</u> strains containing plasmids and having 1/100 volumes, obtained by culturing for 12 hours, were added to 10 ml of LB broth containing 50 μ g/ml

of ampicillin. The <u>E.coli</u> strains were cultured for 2 hours at 37°C, subsequently an expression-inducible reagent IPTG (isopropyl-8-thiogaractopylanocide) was added, to obtain a final concentration of 1 mM, and incubation was continued for 4 hours at 42°C.

The resultant culture medium was separated by centrifugation at 5000 rpm for 10 minutes (Hitachi RP 20) and the cells were collected and washed with 5 ml of M9 salt medium. After resolving the washed cells in 2.5 ml of 20 mM Tris-HCl buffer (pH 7.6) containing 10 mM of EDTA, the cells were destroyed by sonication (60 seconds), the mixture was centrifuged at 10,000 rpm for 10 minutes and the resulting supernatent was used as an enzyme solution for detecting metals.

15 EXAMPLE 6

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Synthesis of coelenterazine and its analogues

Coelenterazine and its analogues were synthesised by the method disclosed by Inoue et al. (Published by Japan Chemical Society, Chemistry Letters, 141-144, 1975) and by the method disclosed by Halt et al. (Biochemistry, 18, (11) 2204-2210, 1979).

EXAMPLE 7

Method for detecting metals

The enzyme solution obtained by the above method was dissolved in 30 mM of Tris-HCl buffer soution containing 10 mM of EDTA, 1 μg of the same type of coelenterazine substrate as the natural type, obtained by the synthesis method disclosed by Inoue et al., and 5 μl of 2-mercapto-ethanol were added to the solution to obtain 1 ml of total volume. The solution was allowed to stand for 1 hour in an ice bath and aequorin was reproduced. Then, the concentration of metals was measured.

Each sample containing CaCl₂, SrCl₂ and SmCl₃
(1.5 ml, 30 mM) was poured into the solution of aequorin
thus reproduced, the solution was transferred to the

reaction cell of a spectrophotometer (Mitchell-Hasting photometer) and the luminescence of the sample was measured (1). Further, 30 mM of CaCl₂ was injected into the cell and the luminescence was measured again (2).

The results are shown in the following table:

Metal ion	(1)	(2)
	 	
Ca ²⁺ Sr ²⁺ Sm ³⁺	8.3	· 0
Sr ²⁺	4.9	Ò
Sm ³⁺	3.6	0
Mn ²⁺	0	8.8
Mg ²⁺	0	8.4
Mg ²⁺ Pb ²⁺ Cd ²⁺	0 .	8.3
ca ²⁺	0	0.9

20 In the table, the units of the numerals are x10⁻¹¹ photo/sec.

It is apparent from the above table that the metal ions Ca^{2+} , Sr^{2+} and Sm^{3+} can be detected and the presence of Cd^{2+} can be also detected because the inhibitory rate of Cd^{2+} is 90%.

Metals were detected using synthesised coelenterazine of the natural type in the above examples. Further, when the metals were detected by using analogues of coelenterazine, similar results were obtained.

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1 CLAIMS:

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- 1. A method for the detection of metals by measuring the luminescence of a metal in the presence of aequorin, characterised in that the aequorin used has been reproduced by adding coelenterazine or an analogue thereof to an enzyme of aequorin (apoaequorin).
- 2. A method as claimed in claim 1, wherein the aequorin enzyme (apoaequorin) is produced in <u>E.coli</u> by a biosynthesis method.
- 3. A method as claimed in claim 2, wherein the biosynthesis method is a recombinant DNA technique and the aequorin enzyme (apoaequorin) is produced by using aequorin cDNA.
- 15 4. A method as claimed in claim 1, 2 or 3, wherein the metal detected is selected from Ca^{2+} , Sr^{2+} and Sm^{3+} , Cd^{2+} and the lanthanoids.
 - 5. A method as claimed in claim 4, wherein the lanthanoids are selected from ${\rm La}^{3+}$, ${\rm Tb}^{3+}$ and ${\rm Yb}^{3+}$.

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EUROPEAN SEARCH REPORT

Application number

EP 87 30 4051

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Category		Ith indication, where appropriate, vant passages	Relevant to claim	CLASSIFICATION OF THE APPLICATION (Int. Cl.4)
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Place of search THE HAGUE Date of completion of the search 03-08-1987			MEYL	Examiner AERTS H.
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EUROPEAN SEARCH REPORT

Application number

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